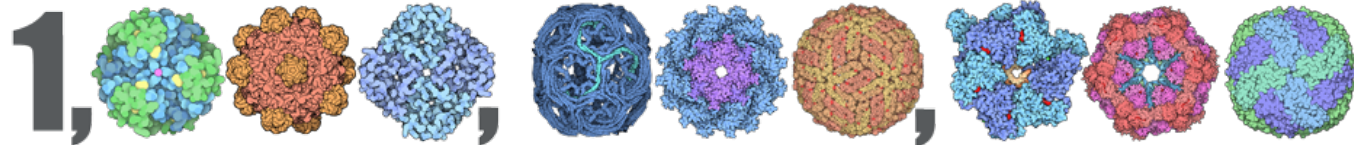
The background image shows the exterior of the San Diego Supercomputer Center building at dusk or night. The building is a modern concrete structure with multiple levels and large glass windows. The entrance features a prominent overhang supported by columns. Small trees are planted along the walkway in front of the building, which is illuminated by streetlights.

Introduction to MMTF-Spark

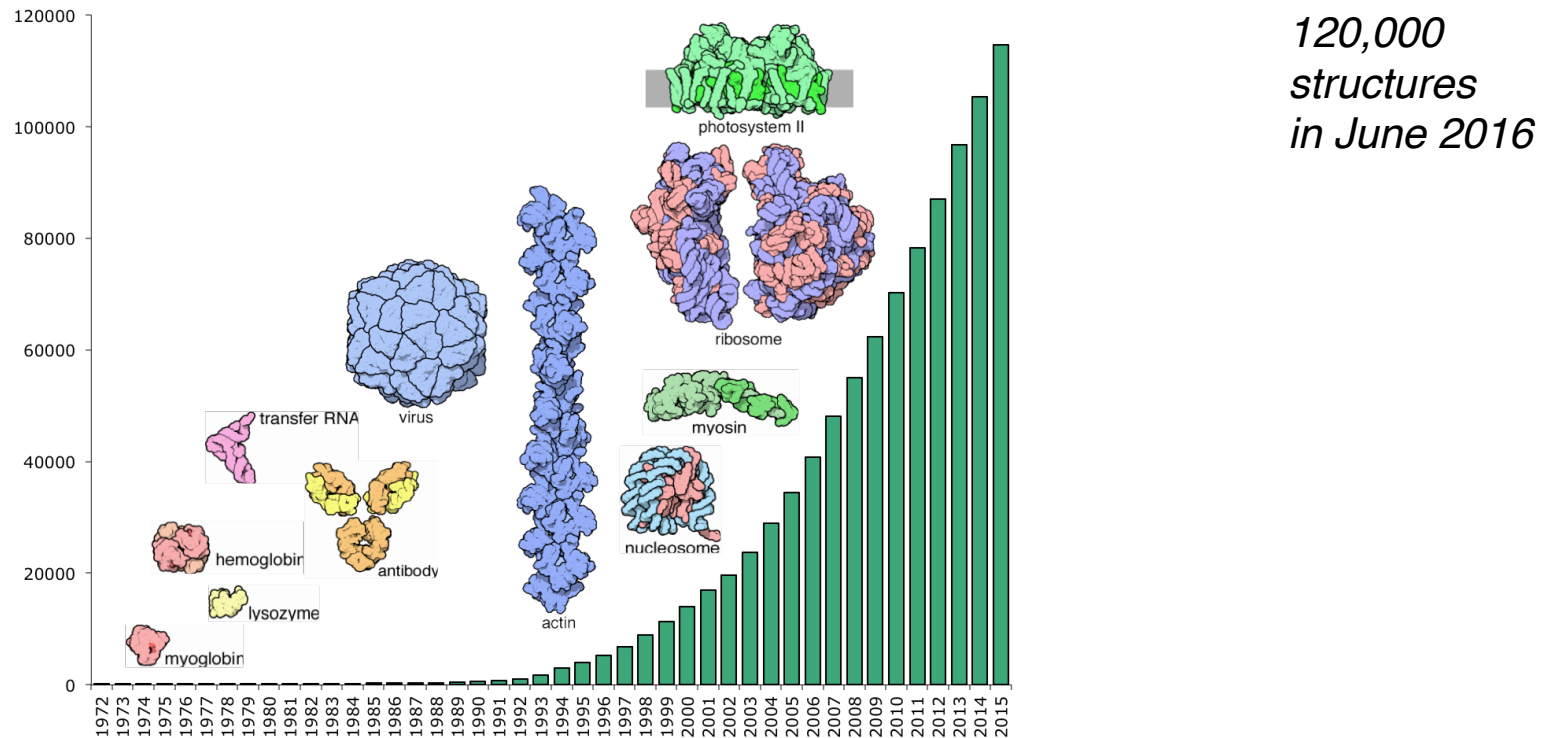
Peter W. Rose, Anthony R. Bradley, Jose M. Duarte,
Antonin Pavelka, Andreas Prlić,
Alexander S. Rose, Yana Valasatava

*Structural Bioinformatics Laboratory
San Diego Supercomputer Center
UC San Diego*

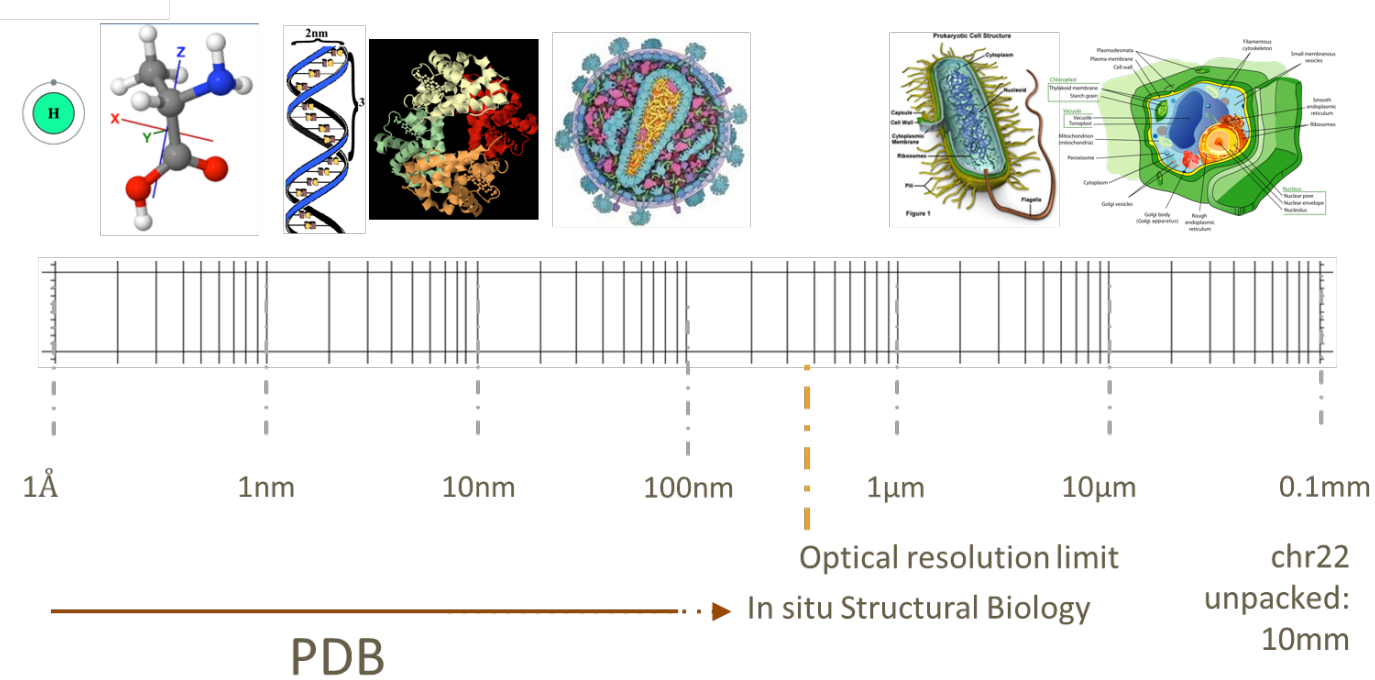
PDB – A Billion Atom Archive



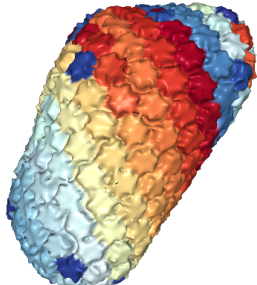
> 1 billion atoms in the asymmetric units



Growing Structure Size and Complexity

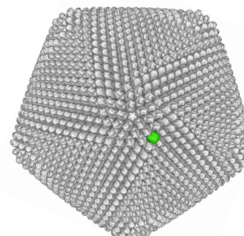


Largest asymmetric structure in PDB



HIV-1 capsid: PDB ID 3J3Q
~2.4M unique atoms

Largest symmetric structure in PDB



Faustovirus major capsid: PDB ID 5J7V
~40M overall atoms

→ Scalability Issues

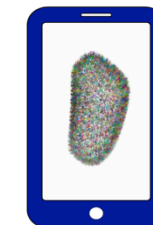
- **Interactive visualization**

- slow network transfer
- slow parsing
- slow rendering



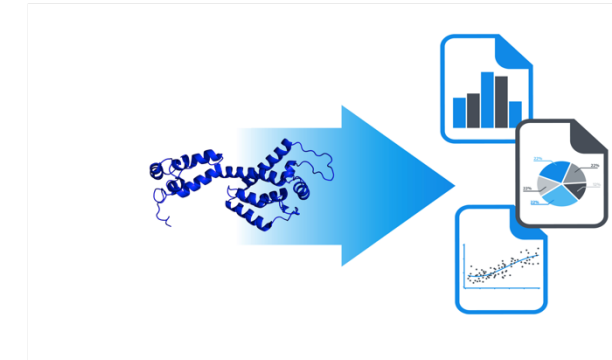
- **Mobile visualization**

- limited bandwidth
- limited memory



- **Large-scale structural analysis**

- slow repeated I/O
- slow repeated parsing

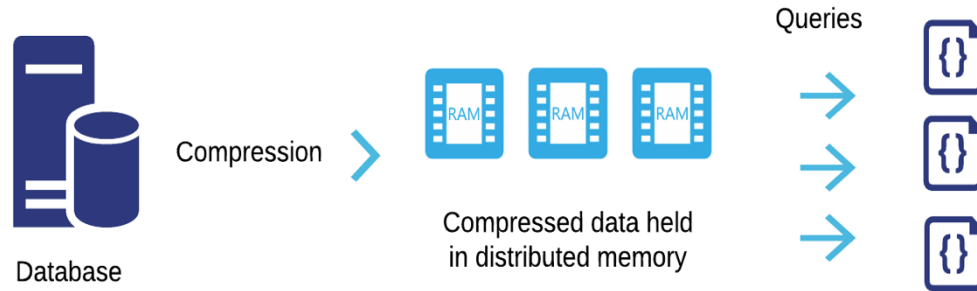


Compressive Structural Bioinformatics

Efficiently store, transmit, and visualize 3D structures of biological macromolecules

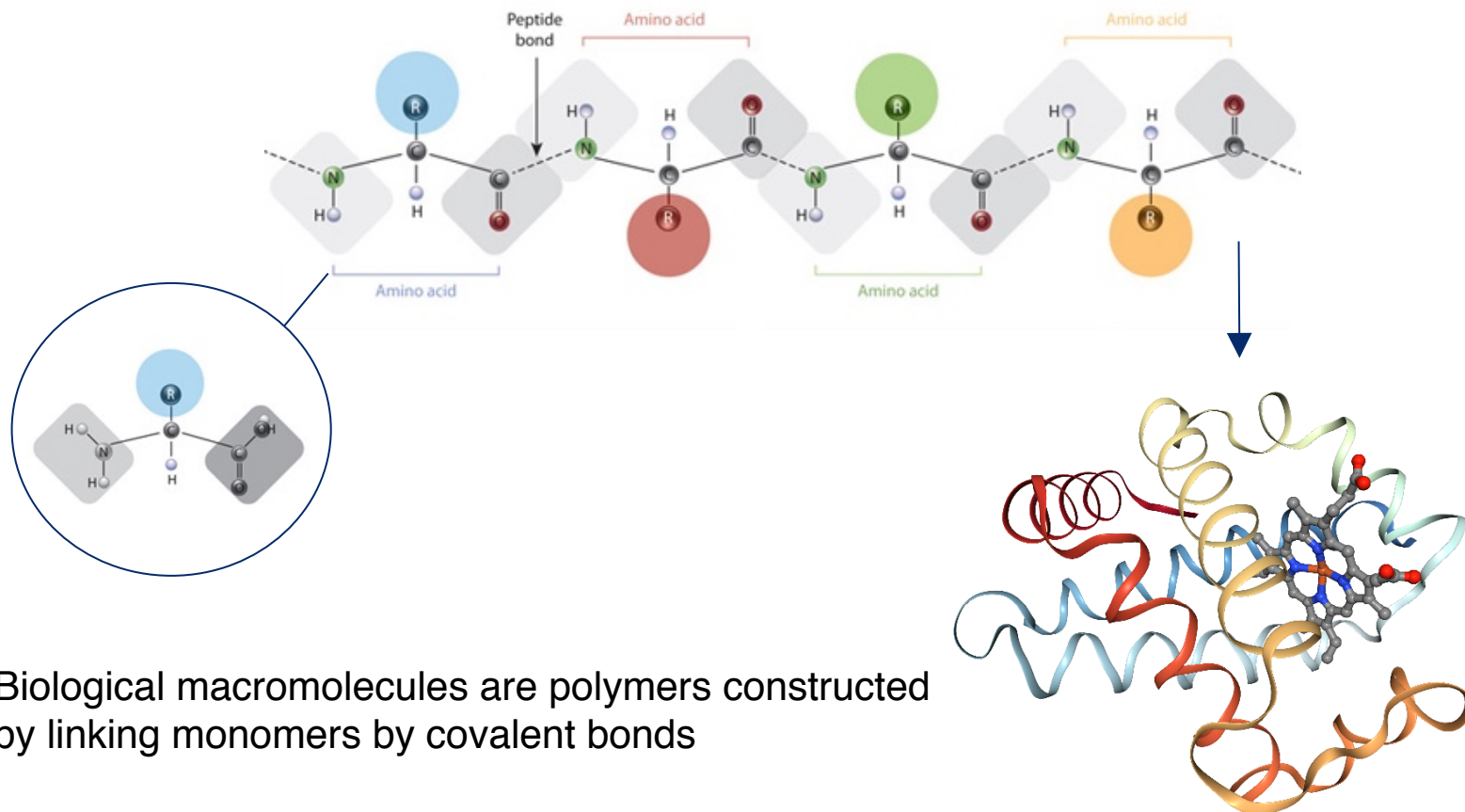


Perform large-scale structural calculations such as geometric queries or structural comparisons over the entire PDB archive held in memory



Macromolecular 3D Structure

Biological macromolecules: proteins, nucleic acids



Biological macromolecules are polymers constructed by linking monomers by covalent bonds

PDBx/mmCIF

```
loop_
_atom_site.group_PDB
_atom_site.id
_atom_site.type_symbol
_atom_site.label_atom_id
_atom_site.label_alt_id
_atom_site.label_comp_id
_atom_site.label_asym_id
_atom_site.label_entity_id
_atom_site.label_seq_id
_atom_site.pdbx_PDB_ins_code
_atom_site.Cartn_x
_atom_site.Cartn_y
_atom_site.Cartn_z
_atom_site.occupancy
_atom_site.B_iso_or_equiv
_atom_site.Cartn_x_esd
_atom_site.Cartn_y_esd
_atom_site.Cartn_z_esd
_atom_site.occupancy_esd
_atom_site.B_iso_or_equiv_esd
_atom_site.pdbx_formal_charge
_atom_site.auth_seq_id
_atom_site.auth_comp_id
_atom_site.auth_asym_id
_atom_site.auth_atom_id
_atom_site.pdbx_PDB_model_num
```

redundant annotations

inefficient representation

repetitive information

ATOM	1	N	N	.	TRP	A	1	5	?	8.519	-0.751	10.738	1.00	13.37	?	?	?	?	?	?	5	TRP	A	N	1
ATOM	2	C	CA	.	TRP	A	1	5	?	7.743	-1.668	11.585	1.00	13.42	?	?	?	?	?	?	5	TRP	A	CA	1
ATOM	3	C	C	.	TRP	A	1	5	?	6.786	-2.502	10.667	1.00	13.47	?	?	?	?	?	?	5	TRP	A	C	1
ATOM	4	O	O	.	TRP	A	1	5	?	6.422	-2.085	9.607	1.00	13.57	?	?	?	?	?	?	5	TRP	A	O	1
ATOM	5	C	CB	.	TRP	A	1	5	?	6.997	-0.917	12.645	1.00	13.34	?	?	?	?	?	?	5	TRP	A	CB	1
ATOM	6	C	CG	.	TRP	A	1	5	?	5.784	-0.209	12.221	1.00	13.40	?	?	?	?	?	?	5	TRP	A	CG	1
ATOM	7	C	CD1	.	TRP	A	1	5	?	5.681	1.084	11.797	1.00	13.29	?	?	?	?	?	?	5	TRP	A	CD1	1
ATOM	8	C	CD2	.	TRP	A	1	5	?	4.417	-0.667	12.221	1.00	13.34	?	?	?	?	?	?	5	TRP	A	CD2	1
ATOM	9	N	NE1	.	TRP	A	1	5	?	4.388	1.418	11.515	1.00	13.30	?	?	?	?	?	?	5	TRP	A	NE1	1
ATOM	10	C	CE2	.	TRP	A	1	5	?	3.588	0.375	11.797	1.00	13.35	?	?	?	?	?	?	5	TRP	A	CE2	1
ATOM	11	C	CE3	.	TRP	A	1	5	?	3.837	-1.877	12.645	1.00	13.39	?	?	?	?	?	?	5	TRP	A	CE3	1
ATOM	12	C	CZ2	.	TRP	A	1	5	?	2.216	0.208	11.656	1.00	13.39	?	?	?	?	?	?	5	TRP	A	CZ2	1
ATOM	13	C	CZ3	.	TRP	A	1	5	?	2.465	-2.043	12.504	1.00	13.33	?	?	?	?	?	?	5	TRP	A	CZ3	1
ATOM	14	C	CH2	.	TRP	A	1	5	?	1.654	-1.001	12.009	1.00	13.34	?	?	?	?	?	?	5	TRP	A	CH2	1



- **MacroMolecular Transmission Format (mmtf.rcsb.org)**
 - Compact
 - fast network transfer, less I/O
 - Fast to parse
 - binary, no string parsing
 - Contains information for structural analysis and visualization
 - covalent bonds and bond orders
 - consistently calculated secondary structure

Data In MMTF File

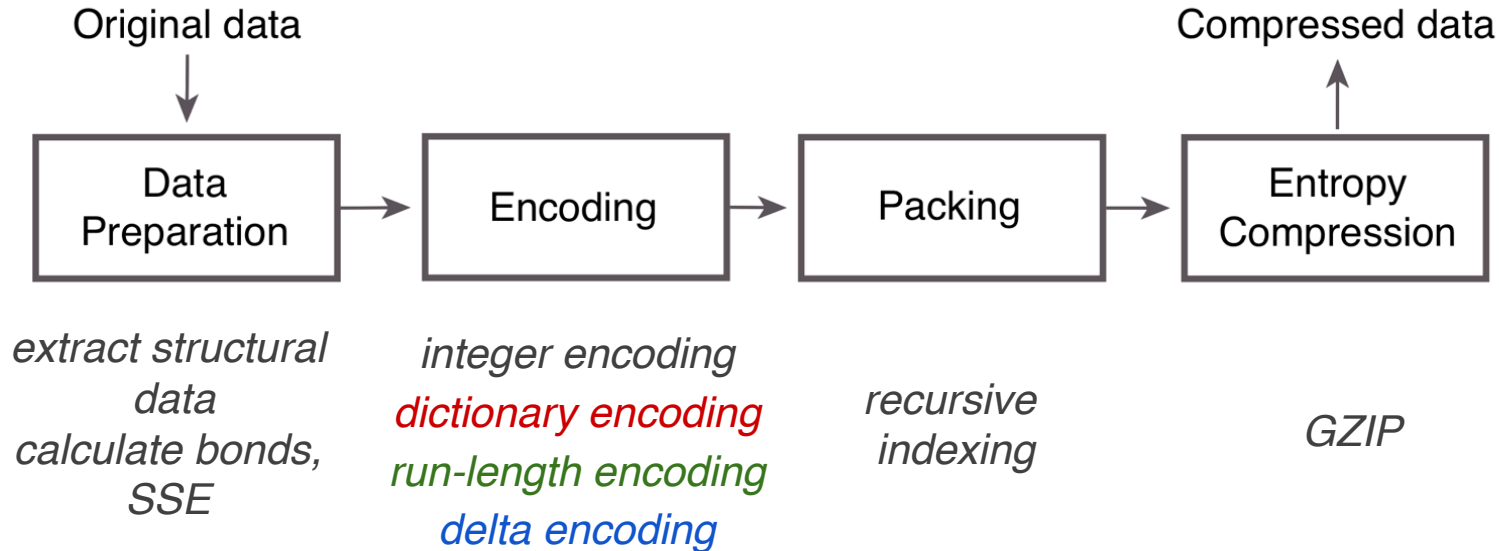
MMTF contains a subset of data used by visualization and structural analysis programs

Data Category	Data Items
Metadata	PDB ID, title, deposition date, release date, experimental method(s)
Crystallographic info	Space group, unit cell, NCS operators, resolution, R_{free} , R_{work}
Primary structure	Polymer sequences
Secondary structure	DSSP secondary structure assignments*
Structural model	Models, chains, groups (residues), atoms, bonds* and bond orders*
Quaternary structure	Biological assembly transformations

* These data items are not available in the PDBx/mmCIF files and are added to MMTF files.

<https://doi.org/10.1371/journal.pcbi.1005575.t001>

MMTF Compression Pipeline



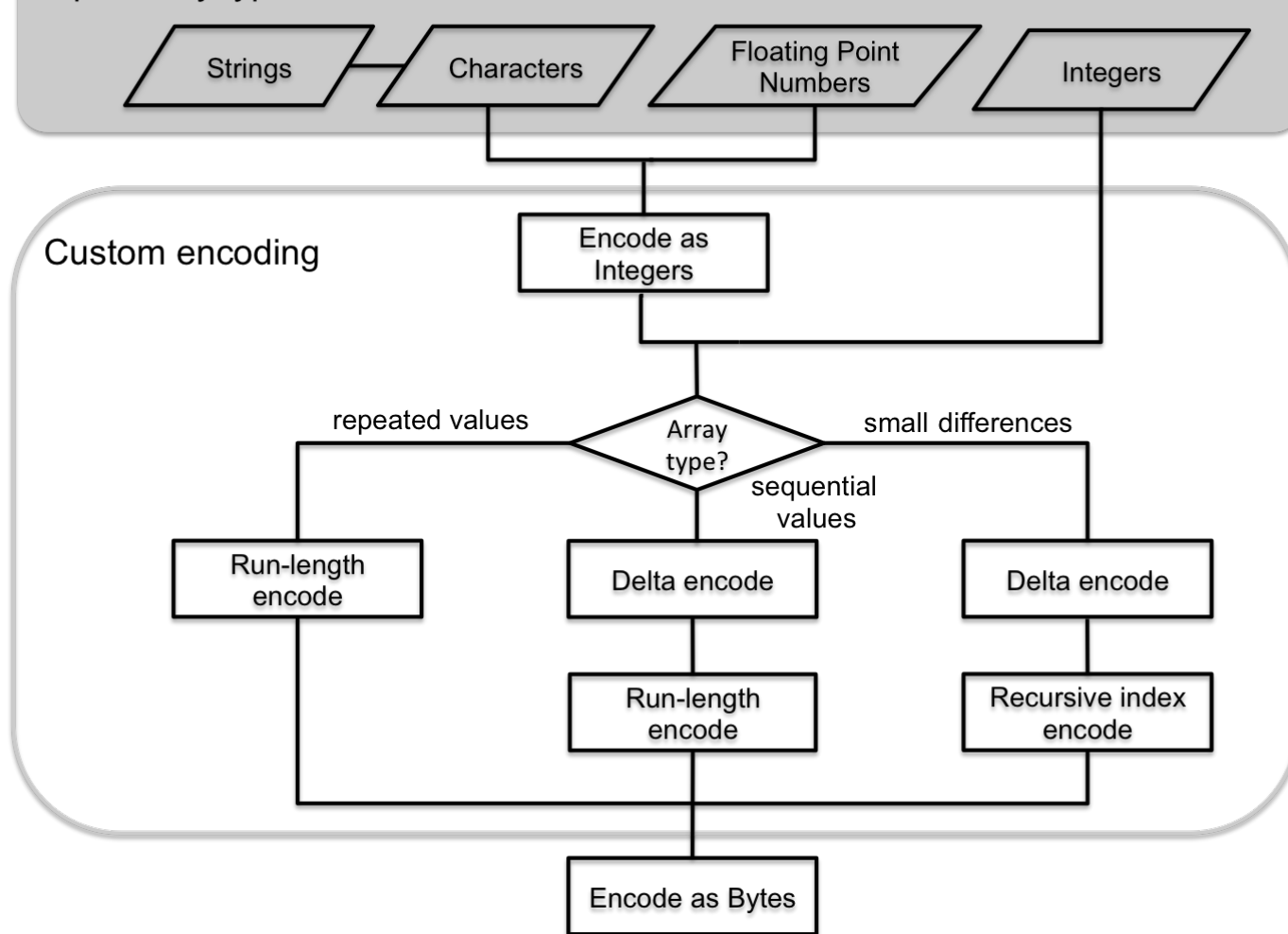
Binary, extensible container format of MMTF

MessagePack

*It's like JSON.
but fast and small.*

Columnar Encoding of Arrays

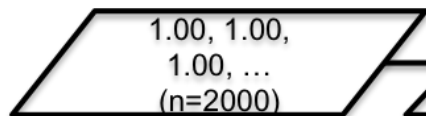
(A) Input array types



Custom Encoding Examples

(B) Repeated values

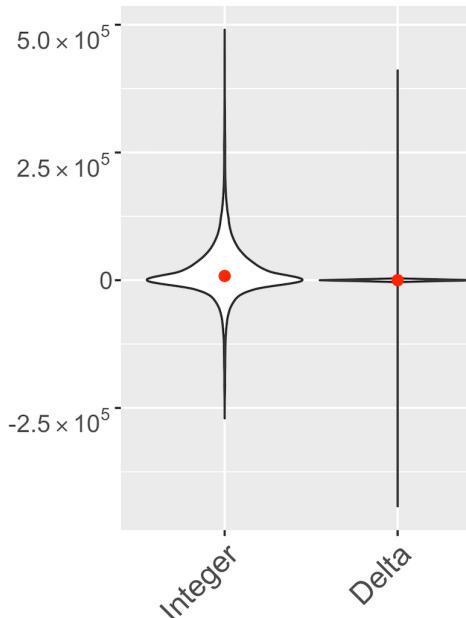
Run-length encoding (e.g., occupancy)



1.00, 1.00,
1.00, ...
(n=2000)

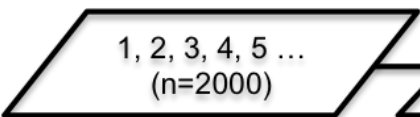
100, 100,
100, ... (n=2000)

100, 2000



(C) Sequential values

Delta and run-length encoding (e.g., serial numbers)



1, 2, 3, 4, 5 ...
(n=2000)

1,1,1,1,1, ...
(n=2000)

1, 2000

(D) Small differences

*Integer, delta, and recursive index encoding
(e.g., xyz-coordinates as 16-bit integers)*



32.867, 34.868,
35.921, ...

32867, 34868,
35921, ...

32867, 2001,
1053, ...

32767, 100,
2001, 1053, ...

Delta encoding reduced the dynamic range of numbers which makes them more compressible

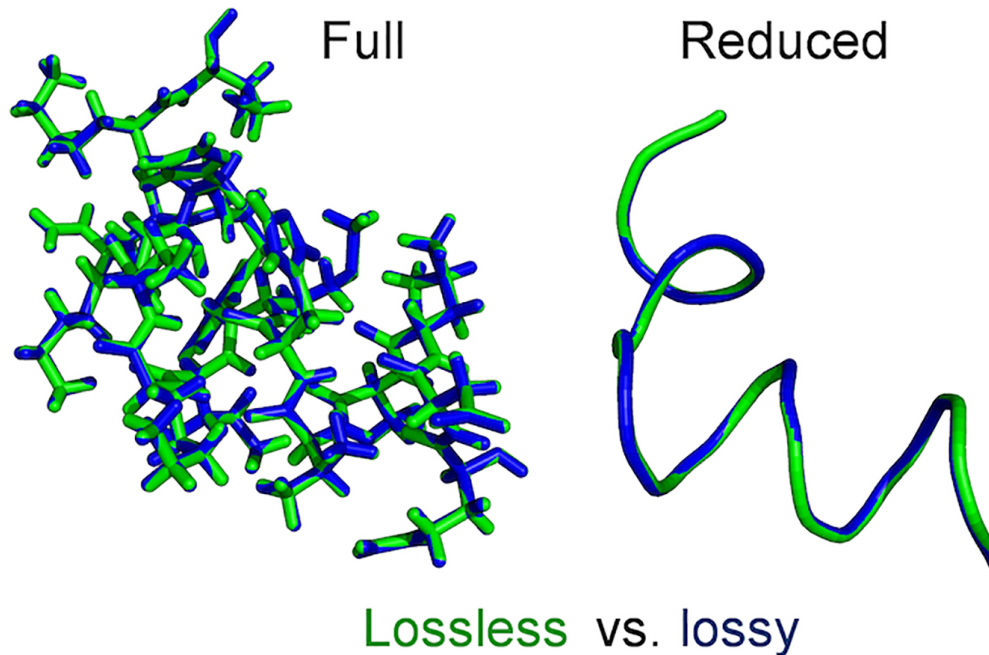
Dictionary Encoding

Unique groups (residues) are stored in a “dictionary”

```
{  
    "groupName": "SER",  
    "singleLetterCode": "S",  
    "chemCompType": "L-PEPTIDE LINKING",  
    "atomNameList": [ "N", "CA", "C", "O", "CB", "OG" ],  
    "elementList": [ "N", "C", "C", "O", "C", "O" ],  
    "formalChargeList": [ 0, 0, 0, 0, 0, 0 ],  
    "bondAtomList": [ 1, 0, 2, 1, 3, 2, 4, 1, 5, 4 ],  
    "bondOrderList": [ 1, 1, 2, 1, 1 ]  
}
```

**Unique entities are stored in a “dictionary”
(e.g., polymer type, polymer sequences)**

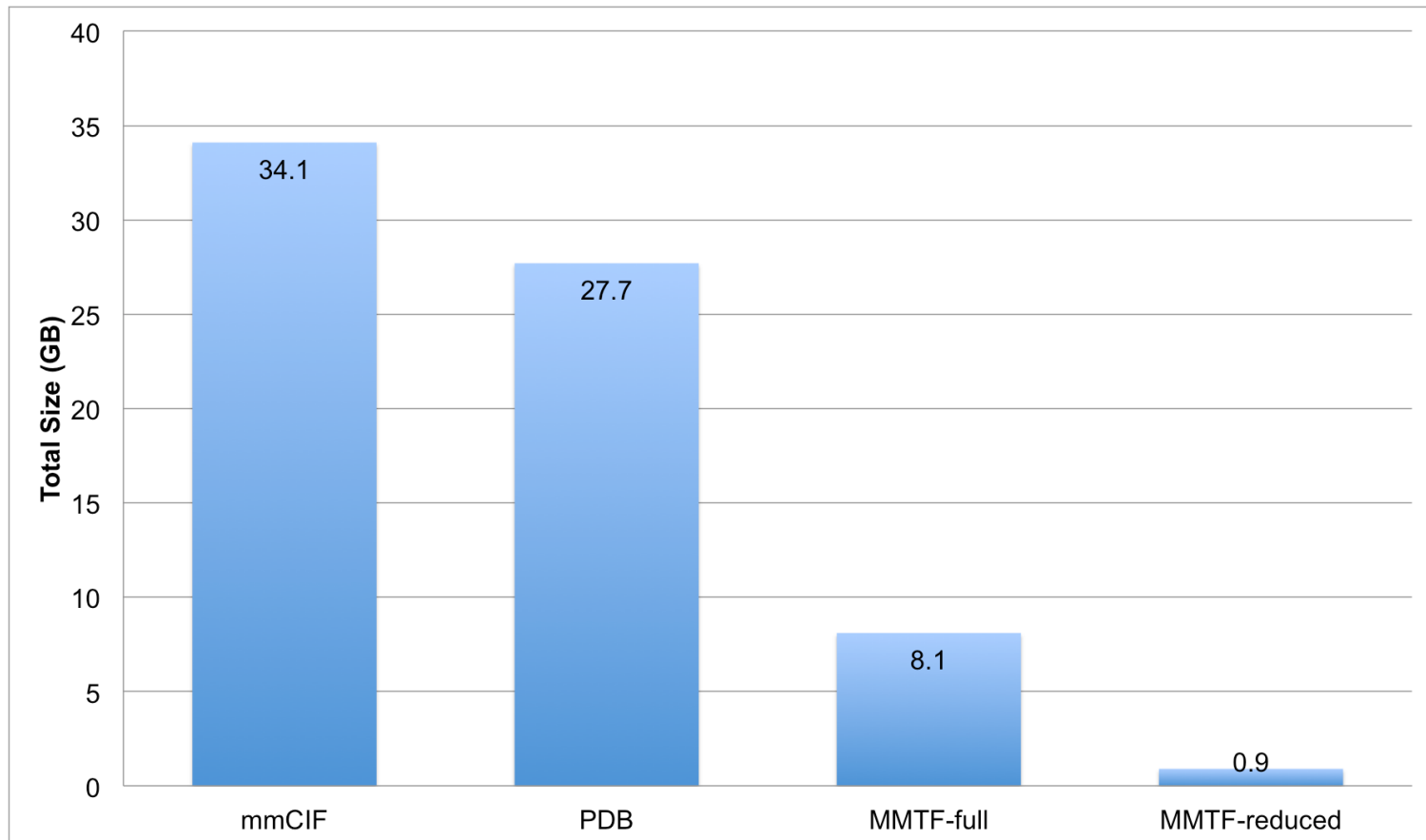
Lossless vs. Lossy Compression



Type	Representation	Coordinate precision (Å)	Temperature factor (Å ²) and Occupancy precision
full	all atoms	0.001	0.01
reduced	C-alpha (polypeptide), P (polynucleotide), all atoms (non-polymers)	0.1	0.1

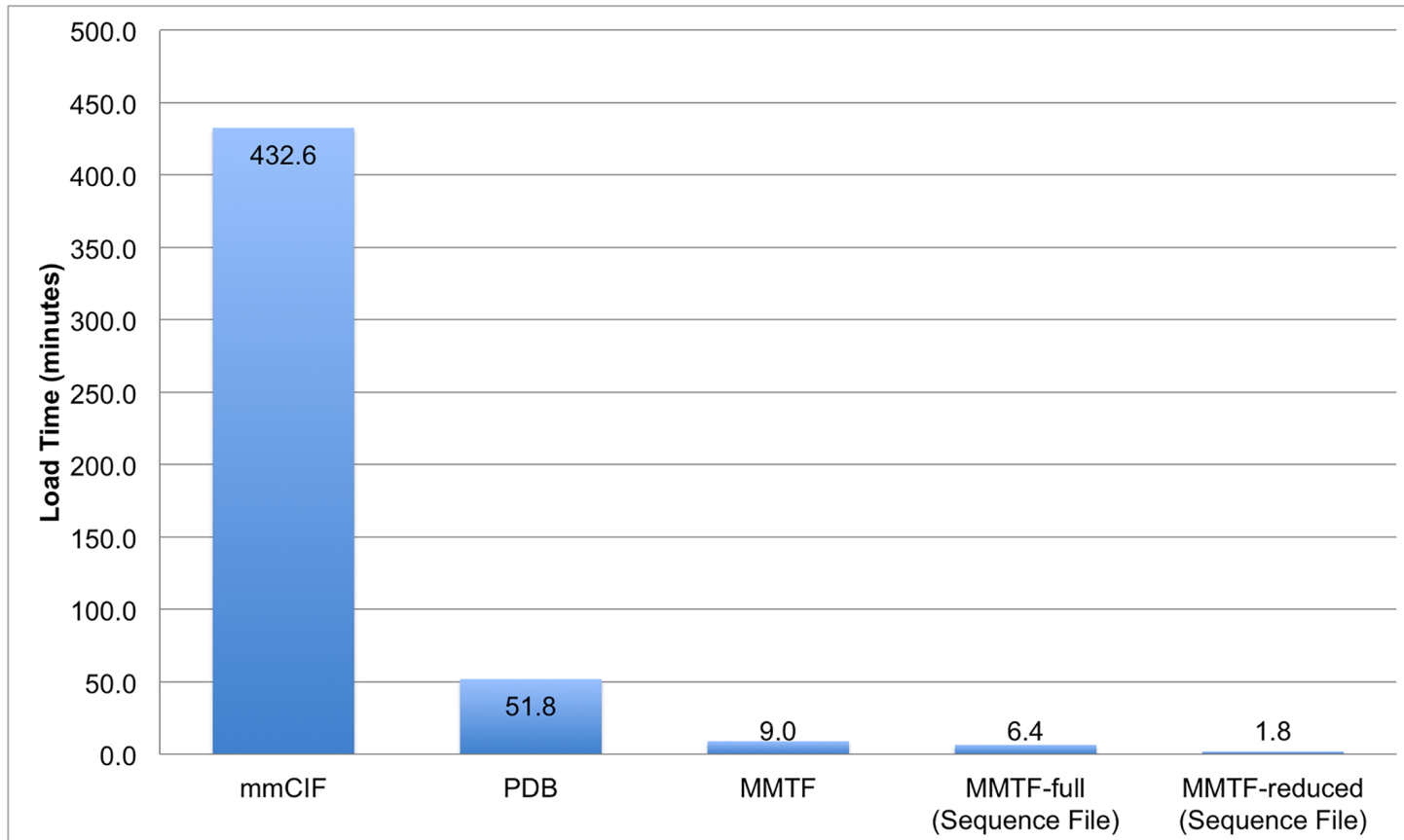
<https://doi.org/10.1371/journal.pcbi.1005575.t002>

File Size Comparison (PDB archive ~127,000 entries)



All file sizes are for the gzip compressed files

File Parsing Speed (PDB archive ~127,000 entries)



Comparison made with BioJava and mmtf-java API (single core)

Download + Parsing time MMTF vs. mmCIF

Time (seconds) to download* 100 large PDB structures from UCSD and parse with JavaScript decoder in Chrome browser



*Note: download times are highly variable and not representative

Community Engagement

- Open source specification
- Open source decoding libraries

Third Party Applications

BioJava,
Jmol

3Dmol.js, iCn3D,
JSmol, NGL

BioPython,
MDAnalysis

PyMol,
ICM Viewer

MMTF API

Java

JavaScript

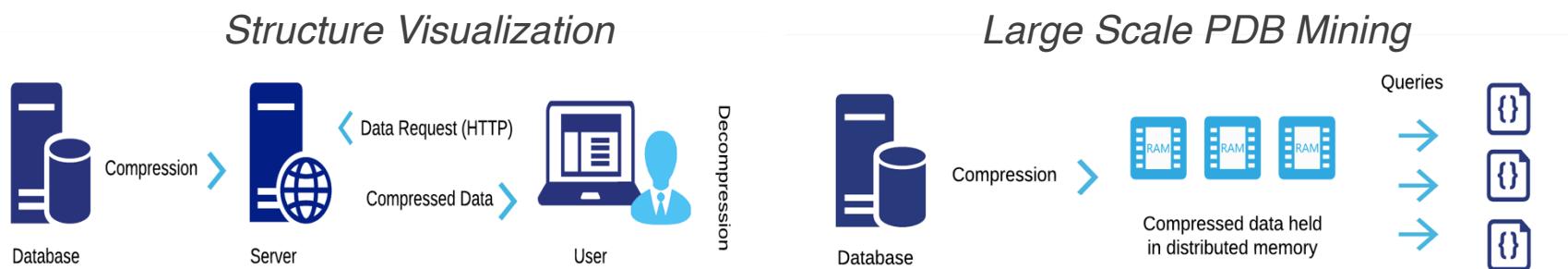
Python

C/C++

MMTF RESTful Web Services

Summary

- **MacroMolecular Transmission Format (MMTF, mmtf.rcsb.org)**
 - Compressed, binary, efficient representation of 3D structures
 - Lossless representation (~4x compression)
 - Lossy, reduced representation (~37x compression)
- **Compressive Structural Bioinformatics**
 - Algorithms, application, and workflows using MMTF
 - 10 to 100+ fold speedup



*Web-based molecular graphics for large complexes (2016)
Web 3D '16, 185-186, DOI: 10.1145/2945292.2945324*

Resources

- **MMTF Website**
 - <http://mmtf.rcsb.org>
- **GitHub Repository**
 - <https://github.com/sbl-sdsc/mmtf-spark>
- **Publications**
 - Bradley AR, et al. (2017) MMTF—An efficient file format for the transmission, visualization, and analysis of macromolecular structures. PLOS Computational Biology 13(6): e1005575. <https://doi.org/10.1371/journal.pcbi.1005575>
 - Valasatava Y, et al. (2017) Towards an efficient compression of 3D coordinates of macromolecular structures. PLOS ONE 12(3): e0174846. <https://doi.org/10.1371/journal.pone.0174846>
 - Rose AS, et al. (2016) Web-based molecular graphics for large complexes. In Proceedings of the 21st International Conference on Web3D Technology (Web3D '16). ACM, New York, NY, USA, 185-186. <https://doi.org/10.1145/2945292.2945324>

Acknowledgements

Funding: NCI/NIH (U01 CA198942)



MMTF Early Adopters

3Dmol.js



BioJava



NGL Viewer

